Application 10/782,435
RuvB Polynucleotides and Uses Thereof

Analysis of SEQ ID NO: 4 performed using the HmmerPfam program in the GCG suite of sequence analysis programs.

Query: 1121SID4 from: 1 to: 455 WPDEF Case 1121 SEQ ID NO: 4 RuvB Scores for sequence family classification (score includes all domains): Score E-value N Model Description _____ 737.6 5.5e-21B 1 TIP49 TIP49 C-terminus ATPase family associated with vario -22.2 0.15 1 Parsed for domains: Model Domain seq-f seq-t hmm-f hmm-t score E-value _____ ~~~~~~ ~~~~ 1/1 117 439 .. 1 326 [] 737.6 5.5e-218 TIP49 Alignments of top-scoring domains: TIP49: domain 1 of 1, from 117 to 439: score 737.6, E = 5.5e-218*->RkAIGvRIKEekeVyEGEVVeleieeaenplgsygktikhgkitLKT R+AIG+RIKE keVyEGEV+el++eeae+ g+y+k i+h++i LKT RRAIGLRIKENKEVYEGEVTELSPEEAESTTGGYAKSISHVIISLKT 163 1121SID4 117 tkmektlkLgpkiyEqLqKEkVeaGDVIyIekntGkVkklGRSdarakdf +k++k+lkL+++iy++L KEkV +GDVIyIe+n+G+Vk++GR+d a+++ 164 VKGTKQLKLDSSIYDALIKEKVAVGDVIYIEANSGAVKRVGRCDSFATEY 213 1121SID4 DlmGeadefVplPkGEVhKrKEvVqtVTLHDlDvaNARpqGglDllSmmg Dl ea+e+Vp+PkGEVhK+KE+Vq+VTLHDlD+aNA+pqGg+D+lS+mg 214 DL--EAEEYVPIPKGEVHKKKEIVQDVTLHDLDAANAQPQGGQDILSLMG 261 1121SID4 aLfspkktEItdelRccINkkVnkwiEcGkAEIVPGVLFIDEVHMLDIEC ++++p+ktEIt++lR+eInk+Vn++i+eG+AE+VPGVLFIDEVHMLDIEC 262 QMMKPRKTEITEKLRQEINKVVNRYIDEGTAELVPGVLFIDEVHMLDIEC 311 1121SID4 FsFLNRALESelAPivilATNRGittIRGTeDiksPHGIPlDLLDR1LII Fs+LNRALBS+1+PivilATNRGi+++RGT D sPHGIP+DLLDRl+II 312 PSYLNRALESPLSPIVILATNRGICNVRGT-DMTSPHGIPVDLLDRLVII 360 1121SID4 tTepYteeEIkkILeiRaqeEgvelseeAldlLtkiGeetSLRYaiqLLt +Te Y + E+ +IL+iRaq+E++++ee l +L +iG +tSLR aiqL+ 361 RTETYGPTEMIQILAIRAQVEEIDMDEESLAYLGEIGQQTSLRHAIQLIS 410 1121SID4 pasilAkkrggkeVeveDVerayeLFlDa<-+ pas+++k++g++++ + D e+++ L 1Da 411 PASVVSKTNGREKICKADLEEVSGLYLDA 439 1121SID4